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Hydroconversion of waxy materials (54)

Hydroconversion of waxy materials, at least a portion of which boils above 700°F (370°C) employs a catalyst containing at least one metal selected from Group VIII, VI and IB supported on a pillared clay. Preferred waxy materials are Fischer-Tropsch and slack waxes.

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Description

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This invention relates to the hydroconversion of waxy materials, especially into distillate materials useful as fuels. More especially, this invention relates to the hydroconversion of waxy materials using a pillared clay catalyst.

Waxy materials such as slack waxes and waxes prepared from hydrocarbon synthesis processes, e.g., the Fischer-Tropsch process, can be upgraded to valuable materials, such as distillate transportation fuels, by hydroconversion processes including hydrocracking coupled with isomerization. The need exists for catalyst with satisfactory conversion and high selectivity to the desired distillate product.

Recent publications regarding the use of pillared clays as supports for hydroconversion processes have been of two minds: pillared clays with a noble metal component are likened to standard noble metal loaded large pore zeolite hydrocracking catalysts where cracking is excessive and selectivity is less than desirable at high conversions, see Doblin, C. et al., Applied Catalysis, 70 (1991) 197-212; while in a separate publication, other authors concluded that insufficient acidity existed in Al-pillared montmorillonites for achieving bifunctional properties, i.e., Molina, R. et al., J. Catal. 148, 304-314 (1994).

In accordance with the present invention, waxes (including Fischer-Tropsch waxes, i.e., materials at least 70% of which boil over 700°F) can be converted to desirable distillate products by treatment with a pillared clay supported metal catalyst through a mild hydrocracking/hydroisomerization process.

Pillared clays are well known in the literature, and their use as hydroconversion catalysts has been reported. See, for example, "Catalysis Today," vol. 2, 2-3 (1988); Pillared Clays, ed. by R. Burch. Nevertheless, the selectivity to desired distillate products where waxy feeds are treated is surprising and unexpected from published literature. Not only are these catalysts highly active for hydroconversion of waxy feeds, the degree of cracking, i.e., gas and naphtha make, is surprisingly lower than would be expected. Thus, the Doblin et al paper, op. cit., disclosed the use of a platinum supported pillared clay for the hydroconversion of octane. Significant cracking was observed. Cracking will be expected to be more severe with waxy feeds, which are more susceptible to cracking than octane. Nevertheless, gas and naphtha make are significantly reduced when waxy feeds are subjected to hydroconversion with the pillared clays described herein, as compared to the use of standard noble metal loaded large pore zeolite hydrocracking catalyst.

Clays useful in preparing pillared clays are swellable clays and may be either naturally occurring or synthetic. Examples of clays that are naturally occurring are: montmorillonite, bentonite, beidellite, nontronite, saponite, and hectorite; and examples of synthetic clays are: fluorohectorite and fluoromicas such as sodium tetrasilicic mica (NaTSM) and synthetic taeniolite.

Pillaring agents, which essentially increase the spacing between the dense clay layers, are generally known and are composed of metal oxides or metal oxide precursors such as alumina, silica, silica-alumina, zirconia, titania and gallium oxide, alone or in combination, preferably zirconia and alumina, more preferably zirconia. After pillaring, the spacing between the layers is known as the gallery height and is generally in the range of about 5-15 Å for pillared clays of this invention. The gallery height is calculated by subtracting the thickness of a clay layer, ~9.6 Å, from the layer repeat distance measured by powder X-ray diffraction.

Suitable clay's useful in this invention are smectites, vermiculites, and micas that may be visualized as a sandwich comprising two outer sheets of silicon tetrahedra and an inner layer of aluminum octahedra (i.e., a 2:1 layered clay). These clays are generally represented by the formula:

$A_x[M_{2-3}T_4O_{10}(Y)_2]$

where M is the octahedral cation, T is the tetrahedral cation, A is the exchangeable interlayer cations $0 \le x \le 1$, and Y is hydroxyl (OH) or fluorine (F) either alone or in combination. The octahedral cations (M) are most commonly cations from the group comprising Al, Mg, Fe, Li; and the tetrahedral cations (T) are most commonly from the group comprising Si, Al, or Fe, either alone or in combination.

A preferred pillared clay and its preparation is described in United States Patent No. 5,248,644 incorporated herein by reference.

In general, however, pillared clay catalysts may be prepared by the following procedure:

The pillared clay or support function of the catalyst is typically prepared by contacting a swellable clay with a solution containing the pillar precursor. Clays that undergo swelling when dispersed in water or another polar solvent are suitable for use in preparing pillared clays. Typical examples of such clays are natural or synthetic clays of the smectite class such as bentonite, montmorillonite, beidellite, saponite, hectorite, and nontronite; vermiculites; and fluoromicas such as taeniolite and sodium tetrasilicic mica. Suitable pillar precursors are compounds of metallic elements such as aluminum, zirconium, silicon, gallium, iron, chromium and the like that form oligimeric species that replace the exchangeable cations between the clay layers when the solution of pillar precursor is contacted with the swellable clay, and furthermore form suitable oxide pillars upon calcination that prop the clay layers apart to create microporous galleries with gallery heights preferably in the range of about 6-14 Å.

Metal functionality for the catalyst can be supplied by metals from Group VIII, noble or non-noble, Group VI, e.g.

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molybdenum, chromium or tungsten, or Group 1B, e.g. copper. Preferred metals are Group VIII metals, particularly the noble metals, most particularly platinum or palladium. Combinations of metals may also be used, for example, Pt/Pd, Ni/W, Co/Mo, Cu/Co/Mo. The amount of metal is present in sufficient quantities to provide hydroconversion activity and may range from about 0.1 wt% to about 20 wt%, depending on the particular metal.

Characteristics of the expanded pillared clay catalyst include surface areas of at least about 100 m²/gm, preferably 100-400 m²/gm, more preferably 250-400 m²/gm; micropore volumes of at least about 0.05 ml/gm, preferably 0.05 - 2.0 ml/gm, more preferably 0.08-1.5 ml/gm.

The dehydrogenation or metal function of the catalyst is added by treating the pillared clay with a suitable metal compound by standard impregnation, ion-exchange, vapor deposition, or other suitable techniques. Metals from Group VIII, VI, and IB can be used individually or in conjunction with one another. Specific examples include Group VIII metals such as Ni, Co, Fe, Pt, Pd, Os, Ru, Ir, and Rh, Group VI metals such as Cr, Mo, and W, and Group IB metals such as Cu. Examples of combinations of these metals that may be used include Pt/Pd, Ni/W, Co/Mo, and Cu/Co/Mo. Preferred are the Group VIII noble metals Pd and Pt.

Waxy feeds useful in this invention are preferably slack waxes or Fischer-Tropsch waxes (preferably from a non-shifting Fischer-Tropsch catalyst), and at least a portion of the feed boils above about 700°F. While 700° F+ feeds are preferred, some 700°F- material may be in the feed and is substantially unaffected by the hydroconversion process. Thus, larger molecules preferentially adsorb in the catalyst and 700°F- materials, in the presence of 700° F+ materials, undergo very little hydroconversion. Thus, hydroconversion is substantially limited to 700°F+ material. The feed is preferably primarily 700° F+, more preferably at least about 70% 700 °F+.

The hydroconversion process is well known in the art and can be effected in the presence of hydrogen over a broad range of elevated temperatures and pressures. For example, temperatures may range from 450 to 750°F, preferably 500 to 700 °F, more preferably 550 to 650°F; while pressures may range from 15-100 atmospheres, preferably 30-70 atmospheres.

Similarly, hydrogen treat rates may vary widely, e.g., 500 to 10,000 SCF/B feed, preferably 1000 to 5000 SCF/B feed. Space velocity may range from 0.5-5 LHSV, preferably from 0.5 - 2.

The hydroconversion is generally carried out at conversions of about 20-80% based on 700°F+ feed. However, there exists a selectivity-conversion trade off. That is, as conversion is increased, selectivity to desired products decreases. Consequently, conversion preferably ranges from about 30-70%, more preferably 40-60% based on 700°F+ feed.

After treatment of the feed with the PILC catalyst, desirable distillate product can be recovered, e.g. by distillation. This product generally boils in the range 250-700°F, and meets required specifications for diesel fuels, i.e., flash point, cloud point, pour point and high cetane number. While the cetane specification is a minimum of 40, the distillate material produced by this process has a cetane number of at least 60, preferably at least 65.

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The following examples will serve to illustrate but not limit this invention.

Example 1 - Preparation of a Zr-Bentonite pillared support

A pillared clay support was prepared from a commercially available montmorillonite clay with zirconyl acetate solution as the pillaring agent using procedures similar to that described in U.S. patent no. 5,248,644. 500 ml zirconyl acetate solution from Magnesium Elektron (ZAA, 22% ZrO₂) was diluted with 2.8 L distilled H₂O and 50.00 g bentonite from American Colloid (Volclay HPM-20) was added. The dispersion was stirred vigorously at room temperature for 3 hr and the solid was isolated via centrifugation and washed with eight 4 L portions of H₂O. The solid product was heated in a fan oven at 120°C for 48 hr. The temperature was then increased at 100 °C/hr to 200 °C, held for 2 hr, increased at 100 °C/hr to 400 °C and held for 2 hr. The yield of Zr-bentonite was 67.3 g, with a layer spacing measured by XRD of 20.5 Å, a BET surface area of 372 m²/g, and a micropore volume of 0.138 ml/g. Elemental analysis via ICP-AES showed 19.5% Zr, 19.2% Si, 7.25% Al, 1.6% Fe, 0.90% Mg, and 0.07% Na.

Example 2

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Catalysts were evaluated for 700°F+ wax conversion activity, selectivity, and product quality under several different sets of processing conditions. The screening was carried out in a small up-flow pilot plant. The catalysts were evaluated at 750 psig H₂, 0.50 LHSV, and with a hydrogen treat rate of 2500 SCF/B. A 10 mL charge of catalyst crushed and screened to 14/35 mesh (Tyler) was employed in all cases. The unit sandbath was equipped with four 0.375" O.D. reactors. Balances were typically collected at 24-72 hour intervals. The reaction temperature was adjusted to provide various 700°F+ conversion levels. The feed was a Fischer-Tropsch wax with the following characteristics:

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	Wt %
I-500	0.70
500-700	20.48
700-1050	62.48
1050+	16.34

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Product distributions were determined by both simulated gas chromatography (i.e., GCD) and 15/5 distillation. Typically, composite samples of 2-4 balance periods were used in the 15/5 distillations. Pour point, cloud point, freeze point, and predicted cetane numbers were obtained on appropriate fractions from the 15/5 distillations using standard ASTM methods. Gas samples were collected in gas collection bombs just prior to the end of each balance and analyzed using mass spectroscopy. 700°F+ conversion was calculated using the following equation:

700 °F+ Conversion = 1 -
$$\frac{700^{\circ}F+ \text{ in product}}{700^{\circ}F+ \text{ in feed}} \times 100$$

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A summary of the catalysts investigated is given in Table 1. Catalyst A was a commercially available material containing 0.50 wt% Pd dispersed on a particulate support containing 70% amorphous silica-alumina with 12% alumina and 30% alumina binder. Catalyst B was a commercially available hydrocracking catalyst containing 0.5 wt% Pd dispersed on a particulate support material containing about 80% ultrastable-Y zeolite and 20 wt% alumina. (Catalyst B was thus similar in nature to the comparative catalyst in Doblin, C., et al., op. cit. which acted similarly as Pt/PILC catalyst in the hydroconversion of octane.) Catalyst C was prepared using the support as described in Example 1 and impregnating to provide 0.5 wt.% Pd thereon..

The metals were impregnated with aqueous solutions of tetraaminepalladium chloride (Strem) using standard incipient wetness techniques followed by drying and calcination.

Table 1

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Catalyst	Composition	Reaction Temperature °F	700 °F+ Conversion
Α	0.5 wt% Pd on amorphous silica-alumina containing 12% alumina and 30% alumina binder	600	52
В	0.5 wt% Pd on composite support with 80% ultrastable-Y	450	49
С	0.5 wt% Pd on zirconia pillared bentonite clay	570	52

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The different catalysts displayed significant differences in wax conversion activity. The most active is the catalyst B while the catalyst C is slightly, but nevertheless significantly, more active than the catalyst A. Activity was determined by the temperature required to achieve ~50% conversion of 700°F+ material in the feed.

Table 2 provides a comparison of the product distributions, jet freeze points, diesel pour points, and cetane ratings

for operations carried out at about 50% 700°F+ conversion.

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5	Catalyst	700 ° F+ Conv .	Product Yields			320- 500 Freeze Point,° C	500- 700 Pour Point, °C	500- 700 Cetan e	
		C1- C4	C5-320	320-500	500-700			*	
	Α	52	1.50	5.29	17.61	38.30	-50.6	-12	66
10	В	49	3.50	13.60	18.03	25.06	-32.0	11	71
	С	52	1.52	5.12	17.46	38.76	-38.5	3	69

Catalyst C had better activity than Catalyst A while producing a similar product slate and had less activity than Catalyst B but produced a markedly superior product slate, yielding more distillate fuel and less gas and less naphtha.

To further investigate the acidity of these catalysts a series of experiments was carried out investigating the isomerization of 2-methylpent-2-ene (2MP2). The 2MP2 isomerization test was carried out as described in G. M. Kramer and G. B. McVicker, Acc. Chem. Res. 19, 78 ff. (1986). The formation rates and rate ratios of the product hexene isomers of this test reaction reflect the relative concentration and strength of the acid sites in the catalyst, respectively. The product hexene isomers formed include 4-methylpent-2-ene (4MP2), trans-3-methylpent-2-ene (3MP2), and 2,3-dimethylbut-2-ene (2,3-DMB2). Formation of 4MP2 requires only a double bond shift, a reaction that can occur on weak acid sites. Formation of 3MP2 takes place through a methyl group shift, which requires a stronger acid site. Finally, formation of the doubly branched 2,3-DMB product requires even stronger acidity. For a given series of catalysts, differences in 3MP2 rates normalized with respect to surface area reflect the density of acid sites possessing strengths sufficient to catalyze skeletal isomerization. Since skeletal isomerization rates generally increase with increasing acid strength, the ratio of the rate of methyl group migration to the rate of double bond shift should increase with increasing acid strength. The use of rate ratios, in lieu of individual conversion rates, is preferable since the difference in acid site populations are normalized.

Table 3 summarizes rates and product distributions for Catalysts A and C. Surprisingly, these results are contrary to the results obtained from the hydroisomerization of a Fischer-Tropsch wax. For example, the higher activity for wax conversion of the PILC catalysts (Catalyst C) relative to amorphous silica-alumina (Catalyst A) shown in Table 1 seems to indicate that the PILCs are more acidic. However, the 2MP2 isomerization test shows the opposite effect. In fact, Catalyst A contains more of the stronger acid sites as measured by the 2,3DMB2/4MP2 ratio as well as more of the weaker acid sites as measured by the 3MP2/4MP2 ratio.

Table 3

2-Methylpent-2-ene Isomerization				
Catalyst	Conversion, wt%	3MP2/4MP2	2,3DMB2/3MP2	
Catalyst A	67.2	1.47	0.192	
Catalyst C	51.6	0.135	0.012	

The results of the 2MP2 isomerization test corroborate the findings of Molina et al. supra, regarding acidity of pillared montmorillonites but the utility of these materials as catalysts is demonstrated herein.

Claims

- A process for the hydroconversion of a waxy material, at least a portion of which boils above about 700 F (370°C), which process comprises contacting the waxy material at hydroconversion conditions with a catalyst containing at least one metal selected from Group VIII, Group VI and Group IB metals supported on a pillared clay.
- The process of claim 1, wherein the waxy material is a Fischer-Tropsch wax.
 - The process of claim 1 or claim 2, wherein the metal is a Group VIII noble metal.
 - The process of claim 3, wherein the metal is platinum or palladium.

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- 5. The process of any preceding claim, wherein the pillared clay has a gallery height of about 5-15 Å (0.5 to 1.5 nm).
- 6. The process of any preceding claim, wherein the pillared clay is a zirconia-pillared montmorillonite.
- 7. The process of any preceding claim, wherein hydroconversion conditions include temperatures ranging from about 500 to 650 F (260 to 345°C) and pressures ranging from about 15 to 100 atmospheres 1.52 to 10.13 MPa).
 - 8. The process of any preceding claim, wherein the hydroconversion conditions are such as to effect from about 30-70% conversion of the 700 F+ (370°C+) material to 700°F- (370°C-) product.
 - 9. The process of claim 1, wherein a hydroconverted product boiling in the range 250-700 F (120-370°C) and having a cetane number of at least 60 is produced.

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(54) Hydroconversion of waxy materials

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EUROPEAN SEARCH REPORT

Application Number EP 97 10 8359

	DOCUMENTS CONSIDE			-1 :		
Category	Citation of document with in of relevant passa	dication, where appropriate. ages		elevant claim	CLASSIFICATION APPLICATION	
x		NOBEL NV ;BOER MARK TUS CAREL (NL); LEL)		3,4	C10G45/64 B01J29/04	
Α	* claims 1,4,6,7.9- * page 9, line 6 - * page 15, line 7 -		6-9)		
X	EP 0 508 005 A (MOB 14 October 1992 * claims 1,2,4,6 * * page 2, line 25 - * page 3, line 20 -	line 29°*	1,5	5,7,8		
P,X	EP 0 753 563 A (EXX CO) 15 January 1997 * claims 1-3,6,8-10 * page 2, line 36 - * example 3 *	*	ING 1-4	1,7,8		
					TECHNICAL F SEARCHED	FIELDS (Int.Cl.5)
					C10G B01J	-
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_	The present search report has	been drawn up for all claims				
	Place of search	Oate of completion of the sea	rch		Examiner	
	THE HAGUE	30 October 19	98	Zuu	rdeeg, B	
X:par Y:par doo A:tec O:noo	CATEGORY OF CITED DOCUMENTS ticularly relevant if taken alone ticularly relevant if combined with anot rument of the same category hnological background national background the mediate document	E : earliér pat after the fi ther D : document L : document	ent documenting date cited in the cited for other	application reasons	ished on, or	